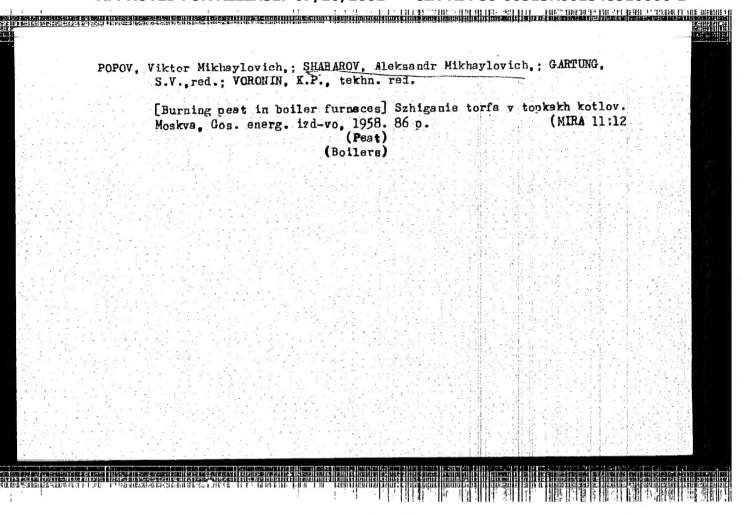


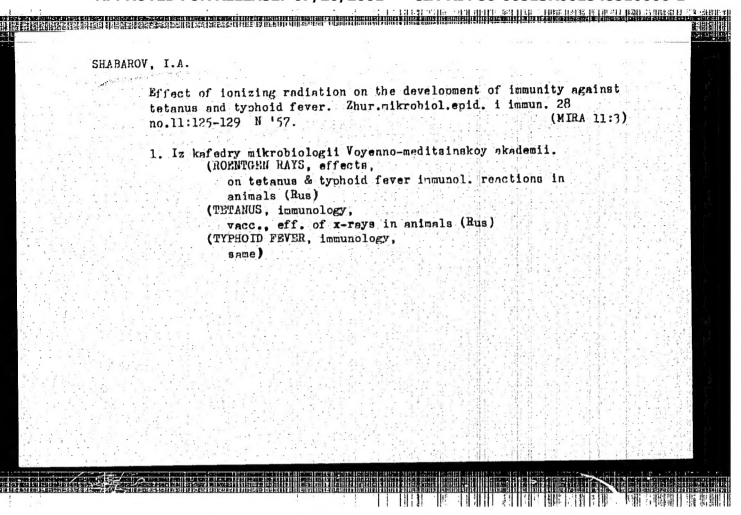
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9. Monthly List of Russian Accessions, Library of Congress, February 1953. Unclassified

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S/032/61/027/001/030/037 B017/B054

AUTHORS:

Bezborod'ko, M. D., Shabarov, L. I., Podol'skiy, Yu. Ya.,

and Vinogradov. G. V.

TITLE:

Device for Testing the Wear Resistance and Antifriction

Properties of Plastic Materials

PERIODICAL:

Zavodskaya laboratoriya, 1961, Vol. 27, No. 1, pp. 104-106

TEXT: A device was developed for testing the wear resistance and antifriction properties of plastic materials. The plastic material is applied to one of the end surfaces of a thin-walled cylindrical tube, and its friction is determined with a metal surface. The moment of friction is determined tensometrically. The resistance of caprone and Fluoroplast-4 to wear by 40 X (40Kh) steel was established by determining the friction coefficients. Results showed that the friction coefficients increased with time, then slowly dropped, and finally remained constant. This course of change can be explained by an increase in temperature of the contact surfaces. There are 3 figures and 3 Soviet references.

Card 1/1

S/191/62/000/005/010/012 B110/B101

15 Daloo

AUTHORS:

Bezborod'ko, M. D., Vinogradov, G. V., Shabarov, L. I.

TTTLE:

Friction and wear of plastics at high contact pressures

PERIODICAL: Plasticheskiye massy, no. 5, 1962, 53-57

TIXT: Friction was studied at point contact of 19.05 mm clameter balls made from: (1) phenol-formaldehyde resin, (2) glass plastic AP-4 (AG-4) on the basis of phenol-formaldehyde resins ($\sim 60\%$ glass fiber of 5-7 μ diameter), (3) phenol-formaldehyde resin with $\sim 60\%$ sulfite cellulose, (4) metal balls from MX-6 (ShKn-6) steel with 62 R_c hardness. Under 5-100 kg loads the following lubricants were used: (1) the nonpolar naphthene paraifin fraction of HMQ-MC-20 (NPF-MS-20) oil, both pure and with 2% by weight admixtures of tributyl phosphite and "khloref 40"; (2) caster oil and molten stearic acid; (3) cumene; (4) water and aqueous solutions of high heat capacity and thermal conductivity, (5) glycols and multivalent alcohols. The minimum load giving rise to irreversible deformations, served as a criterion for lubricant efficiency. Only a slight effect of the lubricants was found for the pair plastic-plastic.

Card 1/2

Friction and wear of plastics at ...

S/191/62/000/005/010/012 B110/B101

For the pair steel-plastic, intensive wear and surface destruction were observed when using the following lubricants: water; aqueous 5% solution of tartaric acid, citric acid, pentaerithrite, trimethylol ethane; phenol melt and alcoholic solution of formaldehyde resin. Corrugation of the smooth surface was found for lubricant LINATUM-201 (TSIATIM-201), Hard MC-20 (Npf MS-20), Npf MS-20 + 2% khloref 40, Npf MS-20 + 2% tributyl phosphite, castor oil, ethylene glycol, diethylene glycol and glycerin. Here glycerin proved best, since the diameter of the depression did not increase and the surface of the plastic balls remained smooth over the total load range, even for 5-6 hr. The anti-wear and anti-friction properties of glycerin for the pair steel-plastic may be ascribed to modification of the steel surface (regeneration of the oxide layers and formation of ferrous glycerates). There are 6 figures and 1 table.

Card 2/2

S/683/62/000/000/008/020. E194/E155

AUTHORS: Bezborod'ko, M.D., Vinogradov, G.V.

Podol'skiy, Yu. Ya., and Shabarov, L. I,

TITLE: Four-ball friction machines and modifications of them

for studying the anti-frictional properties and wear

resistance of plastics

SOURCE: Metody ispytaniya na iznashivaniye; trudy soveshchaniya.

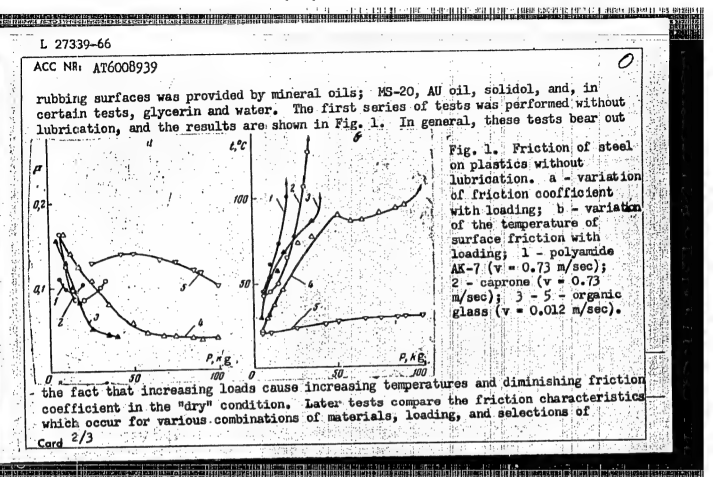
sostoyavshegosya 7-10 dek. 1960. Ed. by

M.M. Khrusbchov. Moscow, Izd-vo AN SSSR, 1962, 81-88

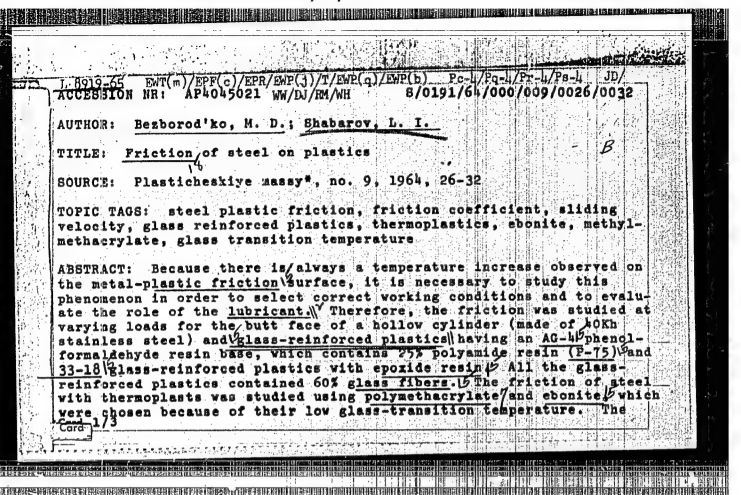
TEXT: Plastic parts are now being extensively used under conditions of sliding with a wide range of loads and speeds. They differ from metals in that their thermal conductivity is low, in that they tend to be of uniform structure throughout, and in that lubricant additives may not act on them in the same way as they do on metals. The sliding properties of plastics should be studied on various materials and with various kinds of lubrication. Four-ball machines can be used, or fixtures adapted for testing two hollow cylindrical specimens in edge contact which can be fitted either in a four-ball machine or in a normal drilling machine. A detailed Card 1/2

5/883/62/000/000/008/020 friction machines and E194/E155 description is given of the fixture for making friction tests between two hollow cylinders. The test specimen being insulated from the machine shaft and from the frame, it is possible to study electrical effects in friction, or the influence of electric current on friction between plastic and metal. The frictional torque is measured by a strain gauge arrangement. In testing, it is important to ensure that the quality of surface finish and contacting of the specimens throughout the surface is uniform. Surface finish is examined with a binocular microscope and plastic specimens may be polished by running-in against a lubricated metal specimen. With the equipment described it was possible to test thermoplastic and thermosetting materials, including reinforced plastics at specific pnessures in the range 2.5 to 300 kg/cm² for flat specimens and up to 8000 kg/cm² in the case of plastic balls at sliding speeds ranging from 0.1 to 20 m/sec. The frictional systems could be operated at temperatures up to 200 °C by circulating a heat-transfer medium. Test results obtained with various combinations of plastica metals and lubricants are described and it is shown that the antifrictional properties and wear-resistance of plastics sliding on metals depend very greatly on the nature of the lubricant. There are 6 figures. Card 2/2

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ACC NR: AT6008939 (A) SOURGE GODE: UR/0000/65/000/0006/0014	
AUTHORS: Bezborod'ko, M. D.; Shabarov, L. I.	
ORG: none	
TITLE: Peculiarities of <u>friction</u> of metals along plastics in <u>lubricating media</u>	
SOURCE: Moscow. Institut mashinovedeniya. Plastmassy v podshipnikakh skol'zheniya; issledovaniya, opyt primeneniya (Plastics in friction bearings; research and experiment in application). Moscow, Tzd-vo Nauka, 1965, 6-14	
TOPIC TAGS: lubricant, friction, friction coefficient, mineral oil, polyamide, material testing, plastic, glycerine, steel, caprone/ hOKh steel, AK-7 polyamide, P6 polyamide, MS-20 mineral oil, AU mineral oil	
ABSTRACT: Experiments on the <u>frictional characteristics</u> of plastics and metals were performed. The object of the investigations was to determine the conditions leading to softening, <u>deformation</u> , and destruction of the plastics. The tests	
were also aimed at the development of an approach to the resolution of the resolutio	
of flicing limits of serviceability of place of a machine which provided friction materials. The experiments were conducted on a machine which provided friction between the planar surface of a steel cylindrical ring on the planar surface of a plastic specimen. Metal specimens were prepared from how the planar surface of a plastic specimen. Metal specimens were prepared from how the planar surface of a plastic specimen.	
plastic specimen. Metal specimens were prepared 1700 and ebonite. Lubrication of were caprone, polyamides P-6 and AK-7, organic glass, and ebonite. Lubrication of Cord 1/3	
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L 8919.-65. ACCESSION NR: AP405021 friction study" was conducted on dry surfaces, in the presence water, glycerine, and high-quality MS-20 oil (naptheneparaffin fracti and at a sliding velocity range of 0.0122 to 1.17 m/sec. The friction was measured at transitions from small to large loads without changing the sample and on a surface area of 78 mm2. The dependence of the coefficient of friction and the temperature on time, the dependence of the friction surface heating and the coefficient of friction on load the dependence of the friction surface heating on the sliding velocity and the effect of the sliding velocity on the heating of steel were studied for glass-reinforced plastics and thermoplastic resins. It was found that at low sliding velocities the lubricant has little influence on the friction between steel and plastics. An increase in the sliding velocity in the absence of lubricants leads to a decrease in the coefficients of friction; brought about as a result of melting of the plastic in the zone of contact with metal. Lowering of friction in the presence of a lubricant is accomplished by a softening of the thermoplastic and chemical modification of the contact surface during friction between the steel and the glass reinforced plastic. In all cases, the decomposition of thermoplastics was a result of heating of the friction surface above the glass-transition temperature and softening of the contact surface. Orig. art. has i

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[Instructions and conventional symbols for making minerel map of the U.S.S.R. on a 1:1000000 scale] Instruktsia i uslovmye obcanacheniia dlia sostavleniia karty poleznykh iskopaemykh SSSH masshtabs 1:1000000. Moskva. Gos.neuchno-tekhn.iad-vo lit-ry po geol. i okhrane nedr. 1955. 16 p. (MIRA 12:10)

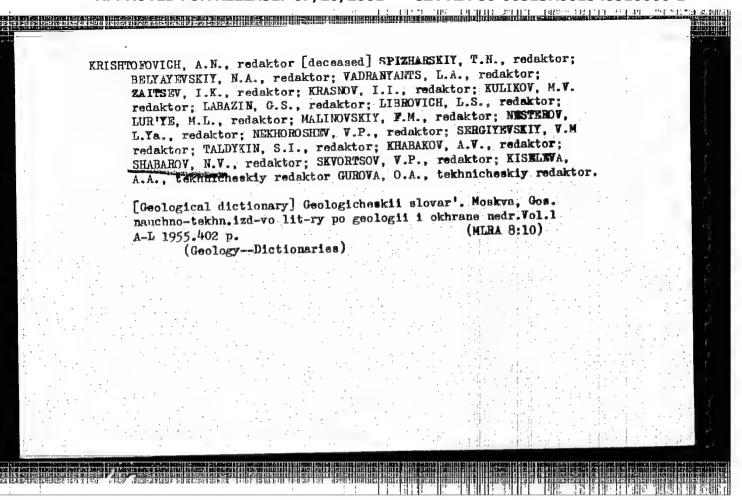
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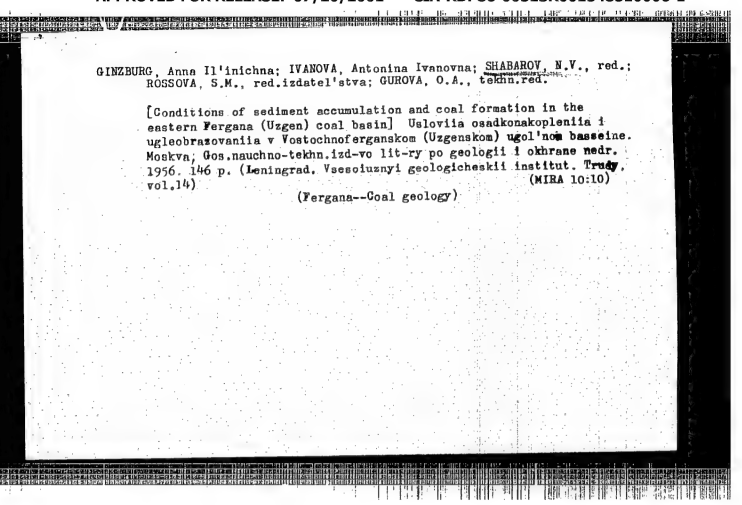
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FOCH, S.G.; GRUSHEVOY, V.G.; DZEVANOVSKIY, Yu.K.; ZORICHEVA, A.I., IVANOV, A.A.; KUREK, N.N.; LIEROVICH, L.S.; MOROZENKO, N.K.; NEKHOROSHEV, V.P.; RUSANOV, B.S.; SPIZHARSKIY, T.N.; SHABAHOV, N.V.; SHATALOV, Ye.T., redaktor; DZEVANOVSKIY, Yu.K.; redaktor; TRASBIKOV; V.I., redaktor; MIRLIN, G.A., redaktor; RUSANOV, B.S., redaktor; SEMENO—VA, M.V., redaktor; GUROVA, O.A., tekhnicheskiy redaktor.

[Instruction for compiling and preparing for publication the state geological map of the U.S.S.R., and the map of the mineral resources of the U.S.S.R.Scale 1:1000000] Instruktsiia po sestavleniiu i podgotovke k izdaniiu gosudarstvennoi geologicheskoi karty SSSR i karty poleznykh iskopaemykh SSSR. Masshtaba 1:1000000. Moskva, Gos. nauchnetekhn. izd-vo lit-ry po geologii i okhrane nedr, 1955. 52 p., tables of symbols, maps [Microfilm] (MIRA 9:6)

1. Russia (1923- U.S.S.R.) Ministerstvo geologii i okhrany nedr. (Geology-Maps)





15-57-8-10394

Translation from: Referativnyy zhurnal, Geologiya, 1957, Nr 8,

p 3 (USSR)

AUTHOR:

Shabarov, N. V.

TITLE:

Vasiliy Ivanovich Yavorskiy (On the Seventieth Anniversary of his Birth) / Vasiliy Ivanovich Yavorskiy (K

vos midesyatiletiyu so dnya rozhdeniya)/

PERIODICAL:

Inform. sb. Vses. n.-i. geol. in-t, 1956, Nr 4,

pp 160-163

ABSTRACT:

V. I. Yavorskiy was born in 1875. He worked as a miner in the coal mines of the Dombrov Basin from 1893 to 1905, at which time he entered upon a course of study at the Petersburg Mining Institute. In 1909, while still a student, he began to work for the Geology Committee. There, under the supervision of L. I. Lutugin, he took part in compiling a detailed geo-

logical map of the Donets Coal Basin. In 1914 Yavorskiy

began his investigations in Kuzbass. In 1927 his

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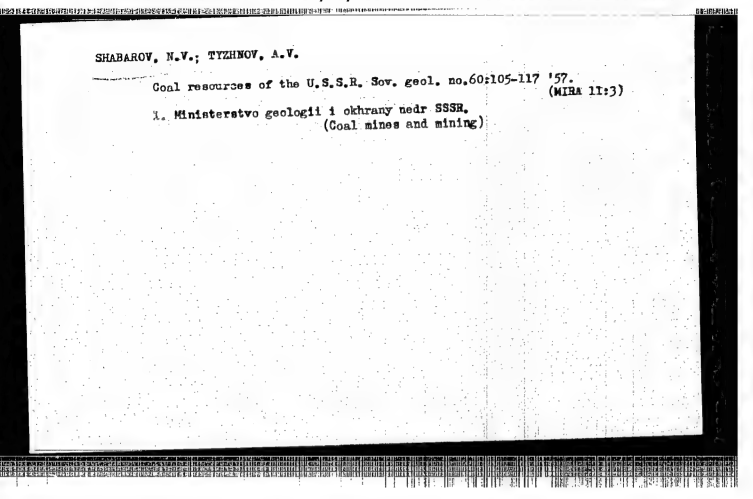
monograph, "The Kuznets Basin", summarized the results

15-57-8-10394

Vasiliy Ivanovich Yavorskiy (Cont.)

of extensive geological studies carried out in Kuzbass by Lutugin's group, much of which work had been done by Yavorskiy himself. Yavorskiy was concerned with questions on the stratigraphy, structural geology, and coal deposits of the Kuznets Basin. He was thus able to play an important part in deciding questions on the Basin development and on its utilization for industry. As a consultant, he selected the sections of the Basin best suited for mining developments. Yavorskiy published more than 160 articles dealing with different problems of coal geology, surveying and industrial utilization of the Kuznets, Donets, and other coal basins of the USSR. He was the author of a series of articles on paleontology and was the sole USSR specialist in stromatology.

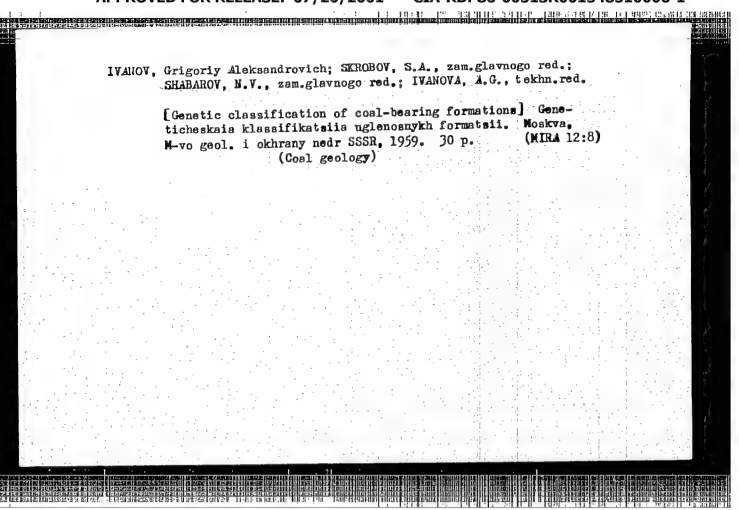
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SHABAROV, N.V., red.; TYZHROV, A.V., red.; VERSTAK, G.V., red.izd-va; AVERKIYEVA, T.A., tekhn.red.

[Reserves of coal and oil shale in the U.S.S.R.; a brief summary of calculations made in 1956] Zapasy uglei i goriucnikh slantsev SSSR; kratkaia svodka rezul'tatov podscheta 1956 g. Pod red. N.V. Shabarova i A.V.Tyzhnova. Moskva, Gos. nauchno-tekhn. izd-vo litry po geol. i okhrane nedr. 1958. 178 p.

1. Russia (1923- U.S.S.R.) Ministerstvo geologii i okhrany nedr. (Coal) (Oil shales)

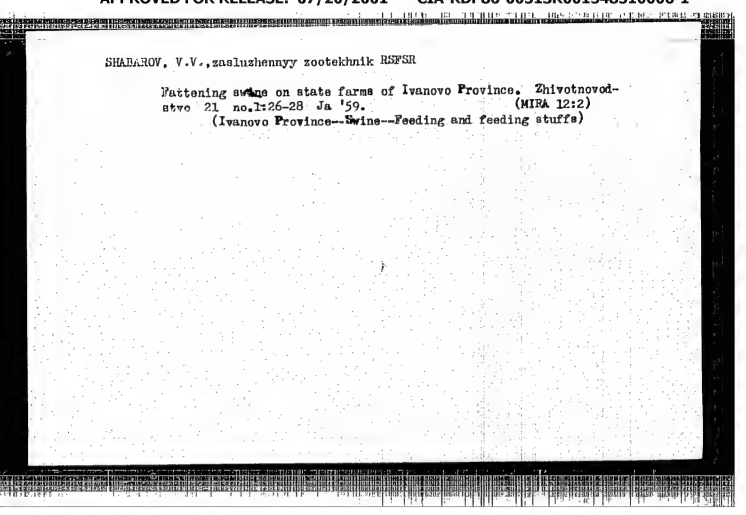


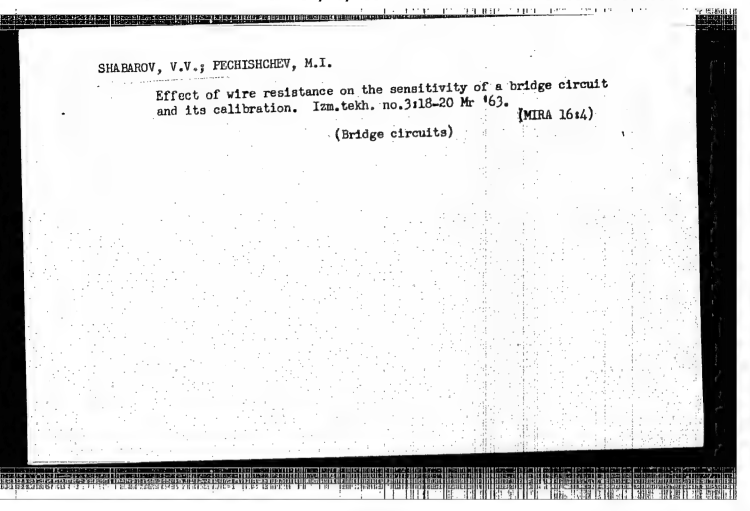
AMMOSOV, I.I., red.; BURTSEV, D.N., red.; GORYUNOV, S.V., red.; GUSEV, A.I., red.; KOROTKOV, G.V., red.; KOTLUKOV, V.A., red.; KUZNETSOV, I.A., red.; MIRONOV, K.V., red.; MOLCHANOV, I.I., red.; NEKIPELOV, V.Ye., red.; PONOMAREV, T.N., red.; POPOV, V.P., red.; PROKHOROV, S.P., red.; SKROBOV, S.A., red.; TYZHNOV, A.V., red.; SHABAROV, N.V., red.; YAVORSKIY, V.I., red.; BOBRYSHEV, A.T., red. toma; VINOGRADOV, B.G., red. toma; VOLKOV, K.Yu., zam. red. toma; LUGOVOY, G.I., zam. red. toma; GGARKOV, V.S., red. toma; SIMONOV, A.V., red. toma; IZRAILEVA, G.A., red.izd-va; IVANOVA, A.G., tekhn. red.

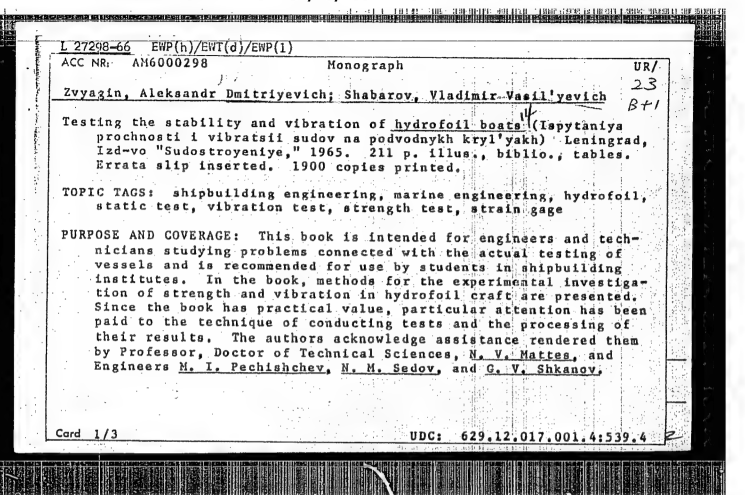
[Geology of coal and combustible shale deposits in the U.S.S.R.] Geologiia mestorozhdenii uglia i goriuchikh slantsev SSSR. Glav.red.I.I.Ammosov i dr. Moskva, Gosgeoltekhizdat. Vol.2. [Moscow Basin and other coal deposits in central and eastern provinces of the European part of the U.S.S.R.] Podmoskovnyi bassein i drugie mestorozhdeniia uglia tsentral nykh i vostochnykh oblastei Evropeiskoi chasti RSFSR. 1962. 569 p. maps. (MIRA 15:9)

1. Russia (1923- U.S.S.R.)Ministerstvo geologii i okhrany nedr.

(Coal geology)

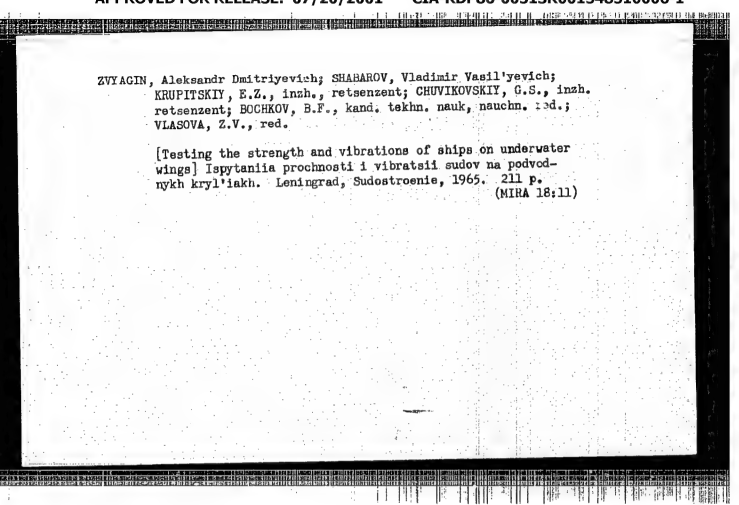






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Ch. I.	Specific feature craft 6	s of strength	and vib	ration	tests for	hydro	foil
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	quantitative ev	aluation 3					
Ch. V.	The effect of sh resistance strai	ipboard test n gage measu	conditio	ns on t 74	he accura	cy of	
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L 27298-66 ACC NR: AM6000298 Ch. VI. The processing of measurement results and the the accuracy -- 108 PART THREE Methods for Strength and Vibration Tests on Hydrofoil Craft Ch. VII. Purpose and problems in testing; devising programs - 129 Ch. VIII. Static strength tests -- 135 Ch. IX. Sea tests -- 146 Ch. X. Vibration tests -- 173 Appendix I. The results of strength and vibration tests performed on the "Vikhr'" ocean-going hydrofoil -- 189 SUB CODE: 13, 14/ SUBM DATE: 03Ju165/ ORIG REF: 048/ OTH REF: 010 Card 3/3



L 10922-67 EWT(d)/EWF(w)/EWF(h)/EMP(1) IJP(c) WM/EK

ACC NR: AR6034806 (N) SOURCE CODE: UR/0398/66/000/008/A024/A024

AUTHOR: Shabarov, V. V.

TITLE: Preparing and conducting motion strength tests of hydrofoil boats

SOURCE: Ref. zh. Vodnyy transport, Abs. 8A141

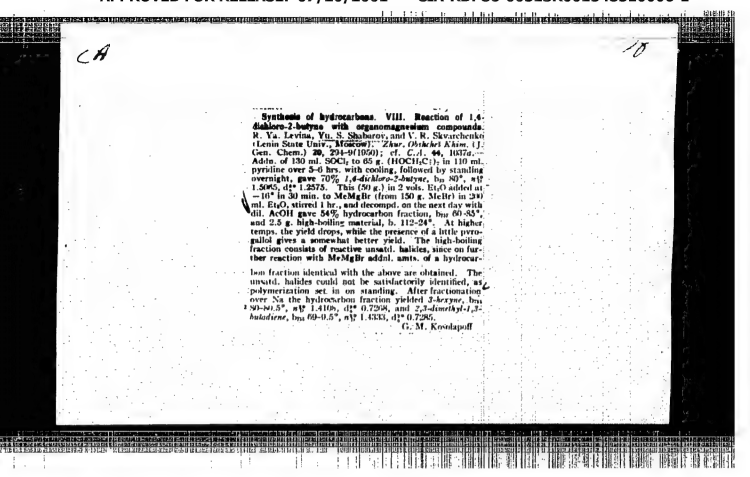
REF SOURCE: Tr. Gor'kovsk. politekhn. in-ta, v. 21, no. 1, 1965, 88-94

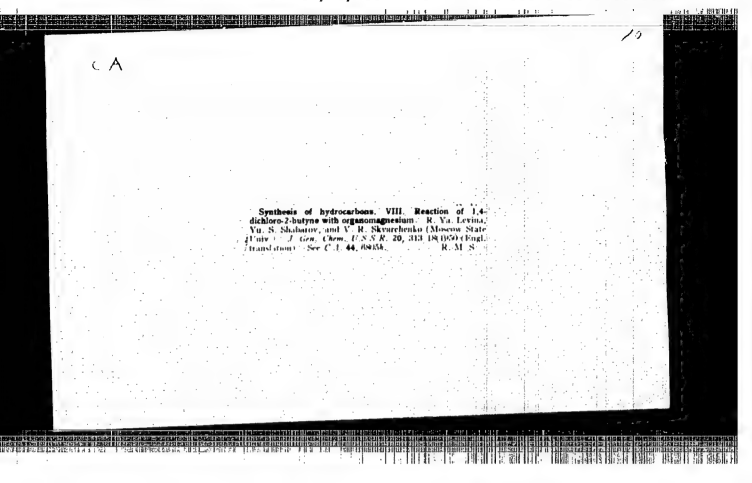
TOPIC TAGS: hydrofoil, motion stability, strength test

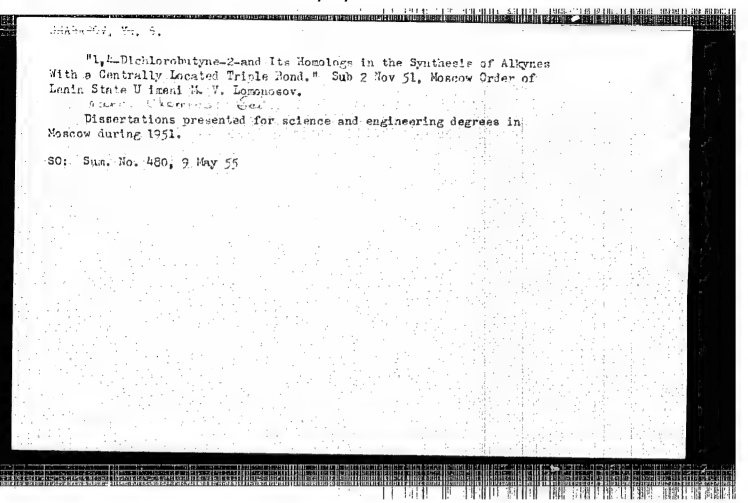
ABSTRACT: A procedure is described for preparing and conducting strength tests of hydrofoil boats. Particular attention is paid to test errors. For measuring the deformation of the hull joints, resistance tensometers, connected by a bridge

hydrofoil boats. Particular attention is paid to test errors. For measuring the deformation of the hull joints, resistance tensometers, connected by a bridge circuit, are used. The tests indicate that during the hydrofoil motion on the sea swell at a speed of 60—100 km/hr, the total duration of stress recording must constitute 300—600 sec at each angle of approach of the hydrofoil motion. Factors are analyzed which affect the operation of the bridge measuring circuit, including atmospheric humidity, temperature of the surrounding medium, coding inaccuracy from pasting the tensometers, nature of the field of deformation, and the length of connecting gears. Bibliography of 3 titles. [Translation of abstract]

Cord 1/1 SUB CODE: 13/ UDC: 629.12.001.4



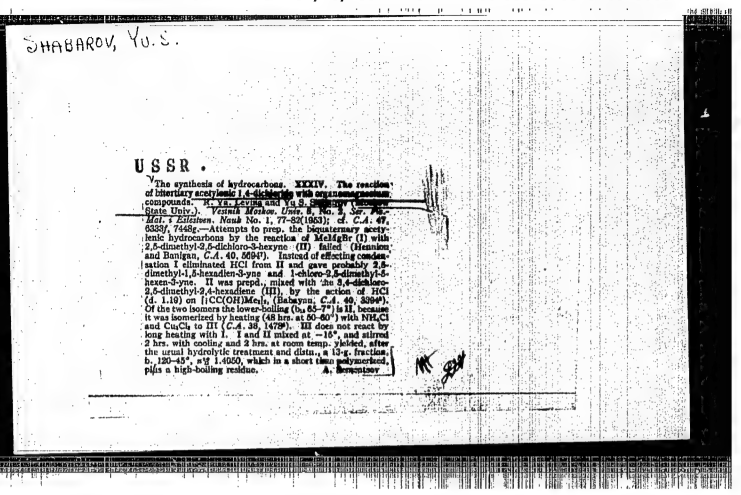


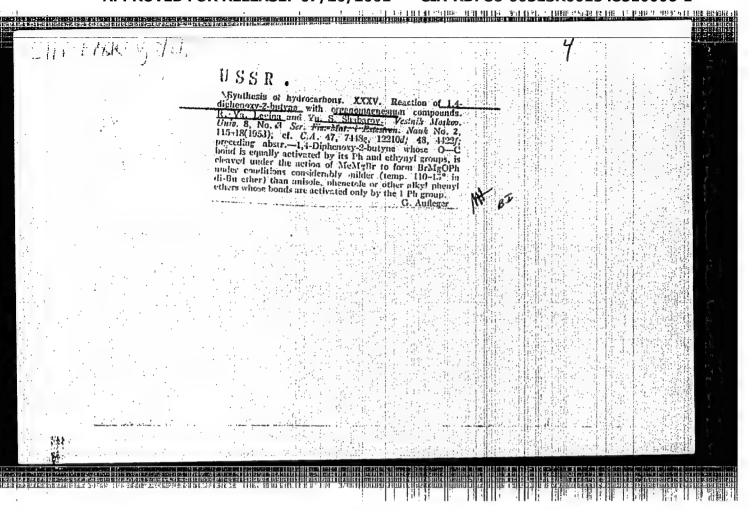


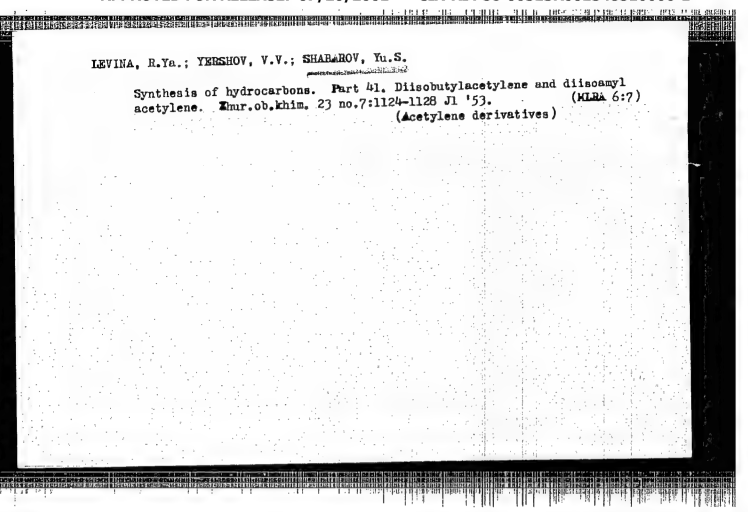
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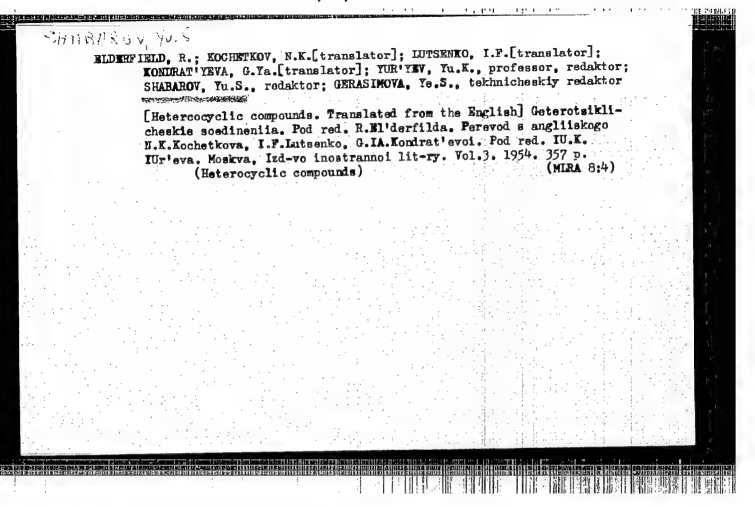
SHABAROV, YU. S.	and dodecyne. ClCH2-C-(ClCH2-C-(cquals C2H5, the reaction to the synthe with a centre Nesmeyanov 26	"Dok Ak Nauk The reaction tylene dichil dichlorbutyn acetylene hy centrally pl	USSR/Chemistry - "The Reaction of nesium Compounds tral Position of Yu. S. Shabarov	
225/13	ne-6. This reaction is shown by: C-C-CH2Cl+2RMgBr+RCH2-C-C-CH2R, whe f, (')H7 and Ch49. The simplicity wi on proceeded shows that it may lend thesis of less readily accessible al tral triple bond. Presented by Acad 26 Mar 52.	"Dok Ak Nauk SSSR" Vol LXXXIV, No 3, pp 509-512 The reaction between methylmagnesiumbromide and ace- tylene dichloride (of the propargyl type, e.g., 1,4- dichlorbutyne-2) was utilized for the synthesis of acetylene hydrocarbons of normal structure with a centrally placed triple bond, like octyne-4, decyne-	Chemistry - Organization 21 May 52 Compounds Reaction of 1,4-Dichlorbutyne-2 With Organomsg- um Compounds Synthesis of Alkines With the Ce Position of the Tripi, Bond, "R. Ya. Levina, Shabarov	

SHIBAROV, Yu. S.	-	232T4	
23274	dichlorides with the chlorine atoms being subscituted by the alkyl group of the Grignard reagent. The secondary dichlorides used as starting The secondary dichlorides used as starting action of thionyl chloride on the corresponding acetylenic gamma-glycols which in turn are easily prepared by the lotsich reaction. In this manner (glycoledichloride-dialkyl-acetylene) many products which are otherwise difficult to prepare synthesized, such as 2,5-dimethylhexyne-3, 3,6-dimethyloctyne-4, and 4,7-dimethyldecyne-5. Presented by Acad A. N. Nesmeyanov 26 Mar 52.	d of Synthesizing Iso-Alkynes in Bond - Di-(Sec-Alkyl)-Acet a, Yu. S. Shabarov SSSR" Vol 84, No 4, pp 709-71 sork, it was shown that Grigos di-primary acetylenes. The samplied to di-sec-acetylenic	USSR/Chemistry - Acetylene Derlyatives 1 Jun 52

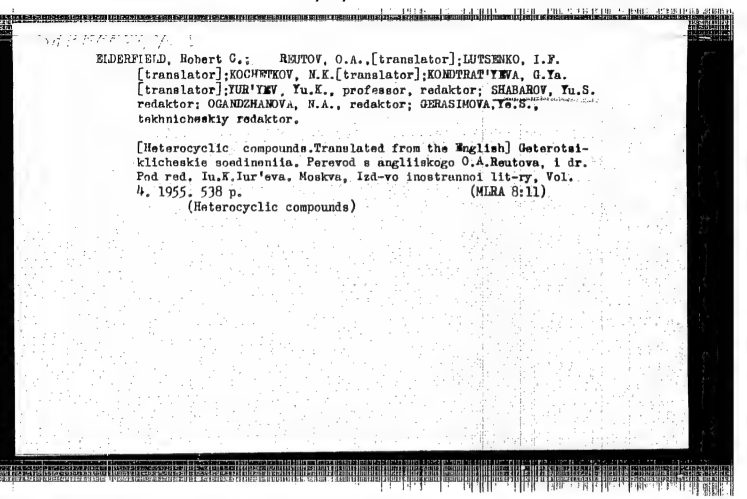


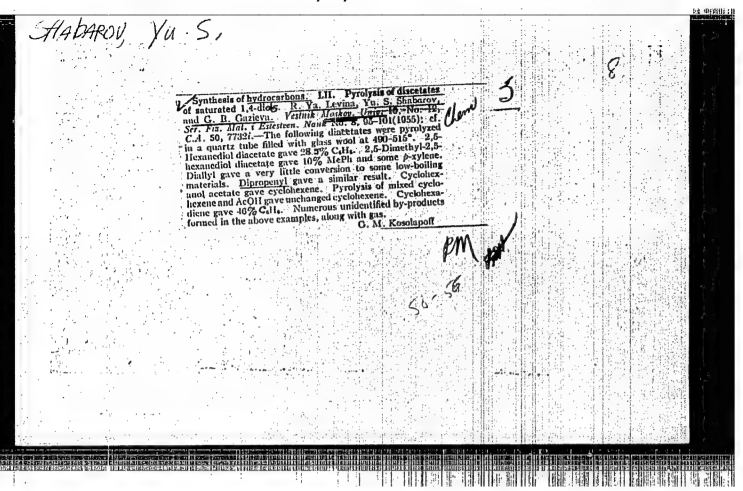


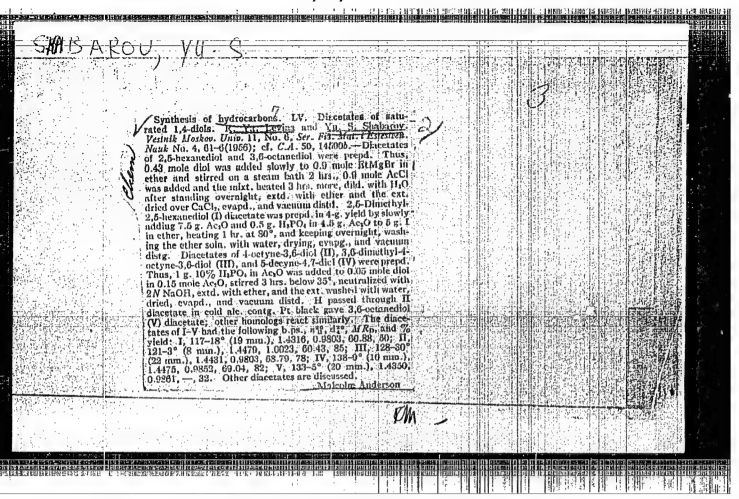


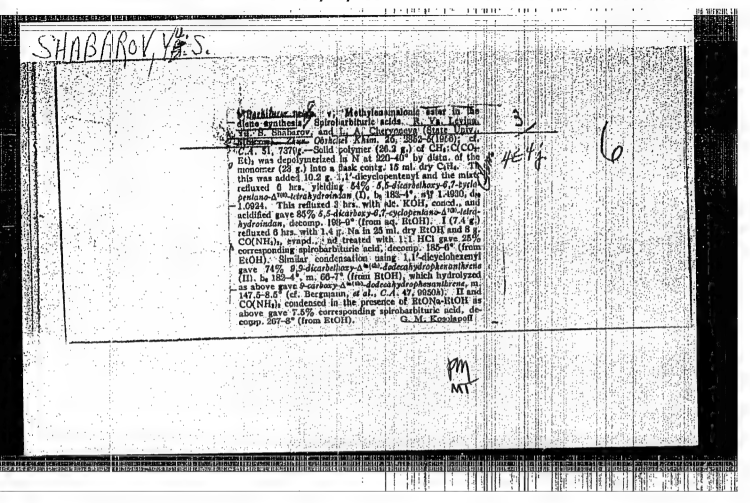


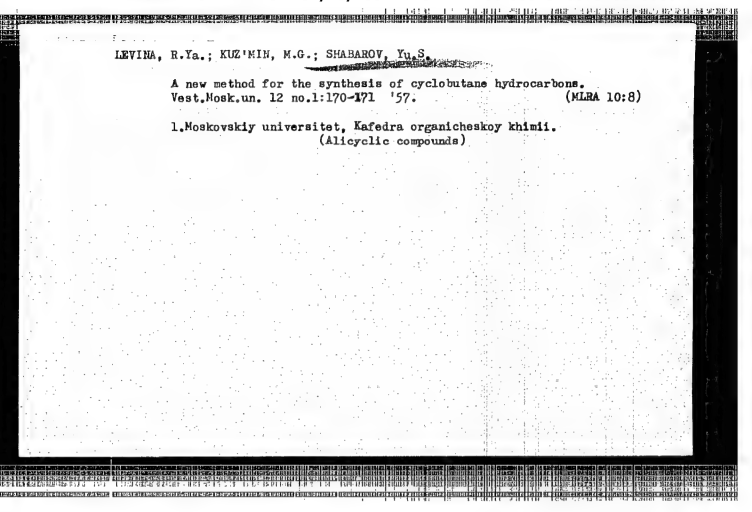
ara Carathera can increase estra a resettine a directina increation de minimistration en interpretarion de man I) El 14 - III - IIII - IIII - IIII - IIII - IIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - IIIII - IIII - III - IIII - III - IIII - III - IIII -. ב.עד איזראמחיי USSR/Chemistry FD-1246 Card 1/1 . Pub. 129-8/25 Author : Levina, R. Ya; Shabarov, Yu. S.; Skvarchenko, V. R. warning and the state of the state of : Hydrocarbon synthesis XLIV. Formation of alkanes resulting from the Title reaction of magnesium-organic compounds with un-saturated 1. 4dichlorides. Vest. Mosk. un., Ser. fizikomat. i yest. nauk, 9, Mo 1, 63-67, Feb 1954 Periodical Established that the formation of paraffin hydrocarbons from the reaction Abstract of a Grignard Reagent with ethylenic and with acetylenic 1, 4-dichlorides is similar. The yield is dependent on the structure of the radical in the Grignard Reagent, the normal type giving a greater yield than those of the iso structure. One table; equations, thirteen references, four foreign. Chair of Organic Chemistry Institution : Submitted July 7, 1953

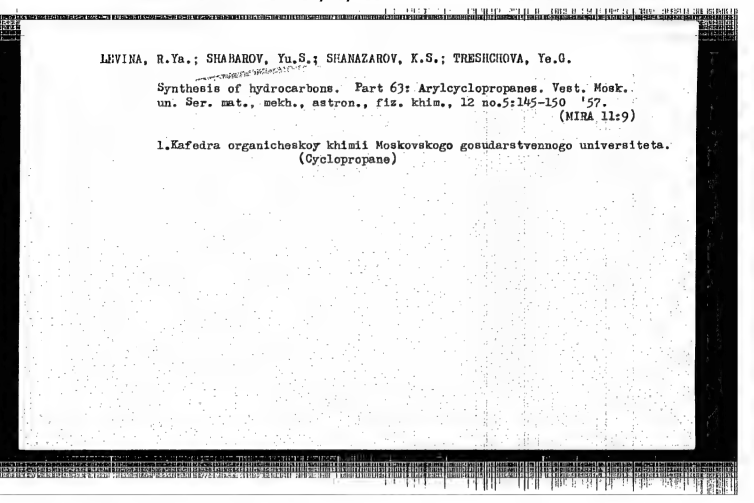












SHINISHREV, YO.S.

AUTHOR: TITLE:

LEVINA, R. Ya., SHABAROV, Yu. S., DAUKSHAS, V.K. and TRachChOVA, Ye.G. 2,4-Dimethylpentadien-1,3 in the Synthesis of Alkanes with Two Quaternary Carbon Atoms Separated by a CH2-Group (Ditertiary

Alkylmethanes).

2,4-dimetilpentadien-1,3 v sinteze alkanov s dvumya chetvertichnymi atomami ugleroda, razdelennymi CH, gruppoy (ditretichnoalkil-

metanov. Russian).

PERIODICAL:

Doklady Akademii Nauk SSSR, 1957, Vol 113, Nr 6, pp 1286 - 1289

(U.S.S.R.)

ABSTRACT:

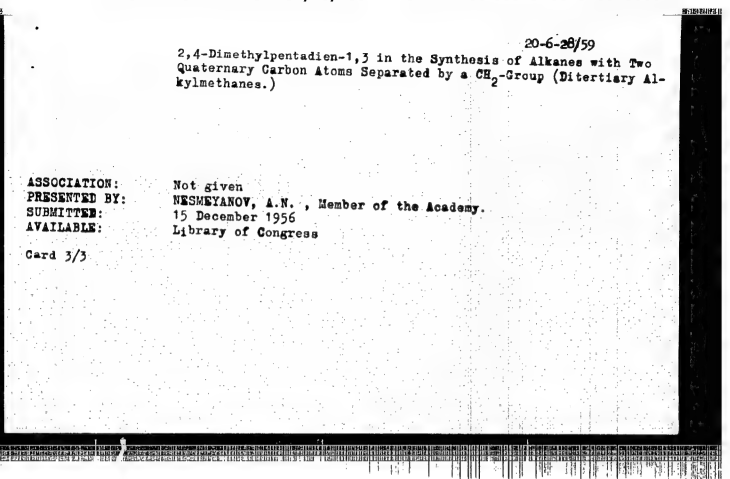
In earlier reports the authors described the method of synthesis of ethylene-hydrocarbons with a quaternary carbon atom. It consists of a reaction between alkylmagnesiumbromides and unsaturated tertiar bromides of the allyle-type; the latter are easily produced by means of hydrobromization of dien-hydrocarbons of a ramitied structure with a conjugated system of double bindings. E.g. hydrobromide of 2,4-dimethylpentadien-1,3 served as initial substance for the synthesis of 2,4-trimethylalkenes. In this paper the authors worked out a new and easy synthe for the production of the rather hard obtainable paraffinhydrocarbons with four quarternary carbon atoms which are separated by act -group, that is of the ditertiary alkylmethanes. Here the same initial substance was used. The reaction mentioned is the first stage of this synthesis, where

20-6-28/59

2,4-Dimethylpentadien-1,3 in the Synthesis of Alvanes with Two Quaternary Carbon Atoms Separated by a CH2-Group (Ditertiary Alkylmethanes).

the yield could be increased from 30 to 45 %. Furthermore, the alkenes which had already a quaternary carbon atom were transformed by the action of hydrochloric acid into saturated tertiary chloride 2-chlorine-2,4-trimethylalkanes (yield 90%). The latter were then introduced into the reaction with magnesiumorganic compounds with sublimate as a catalyzer; the products of this last stage of the synthesis were alkanes with two quaternary carbon atoms separated by a CH2-group. In the case of an interaction with magnesiumorganic compounds the saturated tertiary alkanes split off also hydrochloric acid and, with a yield of 50% form the initial alkanes. The latter were used for alkane-synthesis. The akenes and alkanes were easily separated by destillation. The alkane yield amounted to from 15 - 25 % calculated according to the tertiary chloride introduced into the reaction, and to from 30 - 50 % according to that not utilized. Reactions with yields, methods, and spectroscopic constants are described. (3 tables, 6 Slavic references).

Card 2/3



AUTHORS: Levina, R. Ya, Shabarov, Yu.S., and Daukshas, V.K. SOV/55-58-1-25/33

TITLE: Synthesis of Hydrocarbons. 64. The 2,4 Dimethylpentadien - 1,3 in the Synthesis of the Alkanes With Neighboring Quaternary and

Tertiary Atoms of Carbon (Sintez uglevodorodov. 64. 2,4 - dimetilpentadien - 1,3 v sinteze alkanov s sosednimi chetvertichnym i

tretichnym atomami ugleroda)

PERIODICAL: Vestnik Moskovskogo universiteta, Seriya fiziko-matematicheskikh i

yestestvennykh nauk, 1958, Nr 1, pp 187-191 (USSR)

ABSTRACT: The authors used a method developed some times ago (synthesis of hydrocarbons with quaternary atoms of carbon by reciprocal action of the hydrochloride of 2,4 - dimethylpentadien - 1,3 and the

of the hydrochloride of 2,4 - dimethylpentadien - 1,3 and the organic combinations containing magnesium) in order to obtain the following combinations: 2,4,4,5 - tetramethylheptene - 2; 2,4,4,5 - teramethyloctene - 2 and 2,4,4 - trimethyl - 5 - ethylhepten - 2. By hydrogenation of these stuffs the authors obtained:

2,4,4,5 - tetramethylheptan; 2,4,4,5 - tetramethyloctane;

2,4,4 - trimethyl - 5 ethyl heptan.

There are 10 references, 7 of which are Soviet, and 3 American.

ASSOCIATION: Kafedra organicheskoy khimii (Chair of Organic Chemistry)

SUBMITTED: April 20, 1957

Card 1/1

Cyclopropanes and cyclobutanes. Part 4: Method for purification of arylcyclopropanes, produced by the Kizhner reaction. Vest. Mosk.un. Ser.mat., mekh., astron., fiz., khim. 13 no.5:201-203 58.								
(MIRA 12:4) 1. Kafedra organicheskoy khimii Moskovskogo gosudarstvennogo uni-								
versiteta. (Cyclopropane)								

SOV/79-28-8-11/66 feving, R. Ya., Shabarov, Yu. S., Saukshas, V. E. AU MORS: Synthesis of the Hydrocarbons (Sintez uglevodorodov) TTTLI: LAY. Di-Tert, -Alkylmethane From 2,4-Simethylpentadiene-1,3 (LXV. Di-tret.-alkilmetany iz 2,4-dimetilpentadiyene-1,3) Zhurnal obehchey khimii, 1958, Vol. 28, Hr 8, pp.2049-2055(" r . Our rIn connection with a previous publication (Ref 1) the authors AR-ERACT: present the following results in this paper: a method of synthesizing two alkanes which have quarternaries separated by CH, groups has been worked out using 2,4-dimethylpentadiene-1,3 as the starting material. This is a difficult synthesis, and these two compounds were previously almost unknown. The method consists of 4 reactions: 1) preparation of a tertiary ungoturated allyl chloride by hydrochlorinating the 2,4-11methylpentadiene-1,3; 2) the decomposition of this product with alkyl magnesium bromide to give alkenes with a cuarternary carbon atom (45 - 65 % yield); 3) the hydrochlorination of thie product; 4) reaction of the resulting tertiary chlorine (maximum yield: 90 %) with alkyl magnesium bromide in the

Synthesis of the Hydrocarbons. SOV/79-28-8-11/66 LXV. Di-Tert.-Alkylmethane From 2,4-Dimethylpentadiene-1,3

presence of mercurious chloride. Di-tert.-alkylmethanes of symmetrical and unsymmetrical structure were synthesized, and some of these contained both quarternary and tertiary carbon atoms. The yield was 3 - 20 % celculated on the begin of the tertiary chloride, and 6 - 40 % calculated on the basis of the product of reaction 1). Descriptions are given for the first time of a few tertiary monochlorides which have a quarternary carbon in the chain. By hydrochloring ting the 2,4-dimethylpentadiene-1,3 a mono- and dihydrochloride were also prepared. There are 3 tables and 12 references, 10 of which are Soviet.

ACHOCIATION: Moskovskiy gosudarstvennyy universitet

(Moscow State University)

UBMITTED:

July 1, 1957

Card 2/2

AUTHORS:

SOV/20-121-2-30/53 Levina, R. Ya., Shabarov, Yu. S., Kuzmin, M. G., Vasil'yev,

N. I., Treshchova, Ye. G.

TITLE:

A New Method of the Production of Cyclobutane Hydrocarbons (Novyy metod sinteza tsiklobutanovykh uglevodorodov)

PERIODICAL:

Doklady Akademii nauk SSSR, 1958, Vol. 121, Nr 2, pp. 303 -

ABSTRACT:

Shortly the authors wrote about the possibility of a synthesis as mentioned in the title by means of the decomposition of tetra-hydro-pyridazine (Ref 1). In the present paper they investigate this reaction by means of some examples. The last mentioned initial substances are 6-membered analogs of pyrazolines. In the case of their heating in the presence of caustic potash and platinum they decompose under the separation of nitrogen and a formation of cyclobutane hydrocarbons. It showed that the biradicals III forming as intermediates not only do not cyclize but even cleave under the formation of ethylene hydrocarbons. The quantitative ratio between the aryl-cyclobutane formed and the corresponding styrene can be

SOV/20-121-2-30/53

A New Method of the Production of Cyclobutane Hydrocarbons

classified according to the ratio between the quantity of nitrogen and that of ethylene separated in the decomposition of the initial monoaryl-terra-hydro-pyridazine (IIa in IIb). Thus the authors were the first to succeed in extending the range of application of the classical Kizhner reaction which hitherto has been regarded only of use in the synthesis of cyclopropane hydrocarbons. This way the authors synthetized the hitherto not described p-tolyl cyclobutane and 1,2-diphenyl cyclobutane. In an earlier paper (Ref 3) the authors proved that in phenyl cyclopropane there exists a conjugation between the benzene nucleus and the 3-membered cycle. The comparison of the intensities of some of the most intensive frequencies (characteristic of the benzene ring) in the spectra of the combination dispersion of phenyl cyclobutane with the intensities of corresponding frequencies in the spectra of the propenyl benzene, phenyl cyclopropane on the one hand and alkyl benzenes on the other hand proved that the monosubstituted aromatic hydrocarbons are arranged in a series as follows: propenyl benzene > phenyl cyclopropane > phenyl cyclobutane > isopropyl benzene (Table 1). The same frequencies in the spectrum of p-tolyl cyclobutane

Card 2/3

SOV/20-121-2-30/53

A New Method of the Production of Cyclobutane Hydrocarbons

have far higher intensities than in the spectrum of p-methylethyl benzene. This fact proves the existing conjugation between the benzene ring and the 4-membered nucleus.

There are 1 table and 12 references, 9 of which are Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

PRESENTED: February 27, 1958, by A. N. Nesmeyanov, Member, Academy of Sciences, USSR

SUEMITTED: February 26, 1958

THE PERSON REPORTED FOR THE PROPERTY OF THE PR 680511 5.3300 sov/55-59-3-25/32 5(3) Levina, R. Ya., Shabarov, Yu. S. AUTHOR: TITLE: Cyclopropanes and Cyclobutanes Vestnik Moskovskogo universiteta. Seriya matematiki, PERIODICAL: mekhaniki, astronomii, fiziki, khimii, 1959, Nr 3, pp 201 202 (USSR) The authors report that they recently synthetized several ABSTRACT: acryl cyclopropanes (together with V. K. Potapov) and acryl cyclobutanes (together with M. G. Kuz'min, N. I. Vasil'yev, and S. I. Pokraka) which had not yet been described in publi cations, and (together with Ye. G. Treshchova) determined their Raman spectra as well as their ultraviolet absorption spectra (Table). In an earlier paper, in collaboration with B. A. Kazanskiy et al., they proved the connection of the three-membered ring with the aromatic ring on phenylcyclopropane, tolylcyclopropane, p-anisylcyclopropane, 1,1diphenylcyclopropane, and 1,2-diphenylcyclopropane. In continuation of this work, also a connection of the fourmembered ring with the benzene ring was found together with Card 1/2

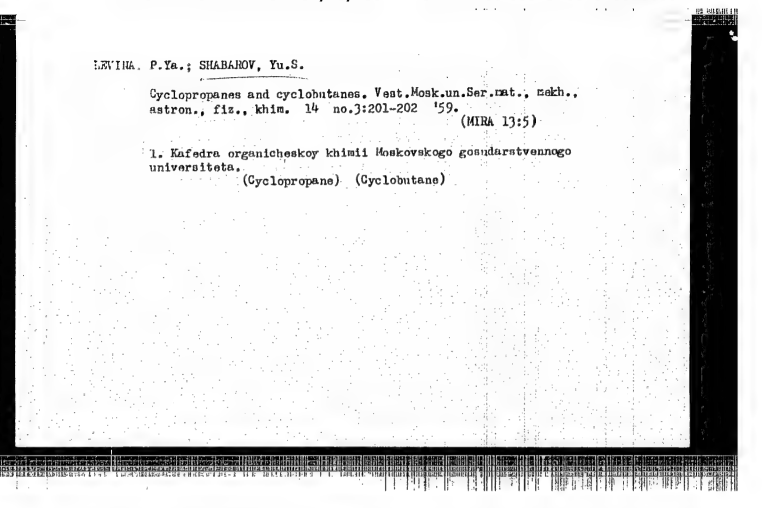
Cyclopropanes and Cyclobutanes

SOV/55-59-5-25/32

M. G. Kuz'min, N. I. Vasil'yev and Ye. G. Treshchova on phenylcyclobutane and p-tolylcyclobutane on the basis of the higher intensities of the characteristic frequencies in the Raman spectrum, which, however, is not so strong as in the case of acrylcyclopropanes. There are 1 table and 4 Soviet references.

ASSOCIATION: Kafedra organicheskoy khimii (Chair of Organic Chemistry)

SUBMITTED: December 24,1958



eration little and the state of the first manifestation and the control of the co SOV/79-29-1-10/74. Levina, R. Ya., Shabarov, Yu. S., AUTHORS: Shanazarova, I. M. Cyclopropanes and Cyclobutanes (Tsiklopropany i tsiklobutany) TITLE: V. Aryl Cyclopropane in the Alkylation Reaction (V. Ariltsiklopropany v reaktsii alkilirovaniya) Zhurnal obshchey khimii, 1959, Vol 29, Nr 1, pp 44-48 (USSR) PERIODICAL: In the present paper the behavior of phenyl cyclopropane was ABSTRACT: investigated in the alkylation reaction of benzene, toluene and anisole as well as that of p-tolyl-and p-anisyl cyclcpropanes in the alkylation reaction of benzene. From the publication data on the alkylation of the aromatic hydrocarbons with alkyl cyclopropanes (Refs 7-12) it follows that AlCl does not cause a preliminary isomerization of cyclopropanes into alkenes (Refs 8,12). The same catalyst was also applied to this case. In the case of alkylation of benzene with phenyl cyclopropane 1,1-diphenyl propane (yield 52%) was obtained, (Scheme 1) the structure of which was confirmed by the reduction of 1,1-diphenyl propene-1 with sodium in butyl alcohol (Schemes 2 and 3). Thus the alkyla-

Cyclopropanes and Cyclobutanes. V. Aryl Cyclopropane in the Alkylation Reaction

SOV/79-29-1-10/74

tion of benzene with phenyl cyclopropane in the presence of AlCl, leads to 1,1-diphenyl propane in the same way as it is the case in the reaction of benzene with styrene which leads to 1,1-diphenyl ethene (Refs 13,14). In the case of the above-mentioned alkylation reaction of benzene, toluene, and anisole an opening of the three-membered ring takes place under the formation of 1-phenyl- 1-aryl propanes (with a yield of 52;61.5;72.5%). It was found that toluene and anisole are alkylated with phenyl cyclopropane in para-position. The alkylation of benzene with p-tolyl-and p-anisyl cyclopropanes proceeds less smoothly than in the case of phenyl cyclopropane as the former polymerize considerably in the presence of AlCl. There are 16 references, 7 of which are Soviet.

ASSOCIATION:

Moskovskiy gosudarstvennyy universitet (Moscow State University)

SUBMITTED:

December 2, 1957

Card 2/2

5 (2) SOV/20-127-1-29/65 Levina, R. Ya., Shabarov, Yu. S., AUTHORS: Kuz'min. M. G. Cyclopropanes and Cyclobutanes (Tsiklopropany i tsiklobutany). TITLE: On the Decomposition of Alkyl-tetrahydropyridazines (O razloznenii alkiltetragidropiridazinov) Doklady Akademii nauk SSSR, 1959, Vol 127, Nr 1, pp 111 - 114 PERHODICAL: (USSR) The authors continue here their investigations of the synthesis ABSTRACT: of aryl-cyclobutanes (Refs 1,2) by the decomposition of aryltetrahydropyridazines and try to use this method for the synthesis of alkyl-cyclobutanes. 3-methyl-tetrahylropyridazine was produced by the hydrolysis of the adduct of azo-dicarboxylic ester with piperylene. The latter contains 2 NH-groups, which indicates that the double bond does not shift towards the nitrogen in the hydrolysis as is the case with the azo-dicarboxylic esters with aryl-butadienes (Refs 2,3), but remains in position 4 (see Scheme). Though the authors failed to produce 3-methyl- Λ^2 -tetrahydropyridazine by hydrolysis, it could be expected that the above-mentioned double bond shifts nevertheless Card 1/4

Cyclopropanes and Cyclobutanes. On the Decomposition SOV/20-127-1-29/65 of Alkyl-tetrahydropyridazines

towards the nitrogen under the difficult conditions of the decomposition and that the compound mentioned is produced. The latter could then be caused to decompose into methyl-cyclobutane and nitrogen (similarly to 3-phenyl- Δ^2 -tetrahydropyzidazine) (Refs 1,2). This failed, however, 3-methyl- Δ^4 -tetrahydropyridazine decomposed with (at 200-250°) or without catalysts (at 300-350°) only into piperylene, nitrogen, and hydrogen (see Scheme). Two reaction ways can be assumed: (a) The biradical I, which is produced as an intermediate, is transformed only into diene. (b) The initial substance decomposes into piperylene and diimide NH - NH which decomposes immediately into N and H. The decomposition of the initial substance into N and piperylene itself in the case of the effect of weak oxidizing agents speaks in favor of the assumption (a). This decomposition confirms the remaining of the double bond in position 4. Similar results were obtained in the decomposition of 3,4,5, 6-bis-cyclopentane- and 3,4,5,6-bis-cyclohexane- Δ^4 -tetrahydropyridazines which contain 2 NH-groups each. They form dicyclopentenyl and dicyclohexenyl besides other reaction pro-

Card 2/4

Cyclopropanes and Cyclobutanes. On the Decomposition SOV/20-127-1-29/65 of Alkyl-tetrahydropyridazines

ducts mentioned here. It was thus found that it is impossible to obtain cyclobutane-hydrocarbons in two stages (hydrolysis and subsequent decomposition) from adducts of the azo-dicarbox-ylic ester with alkyl-butadiene or their cyclic analogs. Therefore, the authors applied another method: they synthesized 3-methyl-tetrahydropyridazine, which, however, turned out to be the isomer of the expected compound with one NH-group. An isomerization of the azo form into a hydrazone form apparently takes place here. Experiments with other substances do not lead to the expected result: either the initial substances were not

transformed, or a resinification took place. Thus, alkyl- Δ^2 -tetrahydropyridazines cannot be decomposed in the presence of catalysts in the case of heating under the precipitation of N and formation of four-membered cyclic hydrocarbons, in contrast

to $aryl-\Delta^2$ -tetrahydropyridazines and pyrazolines. There are 11 references, 8 of which are Soviet.

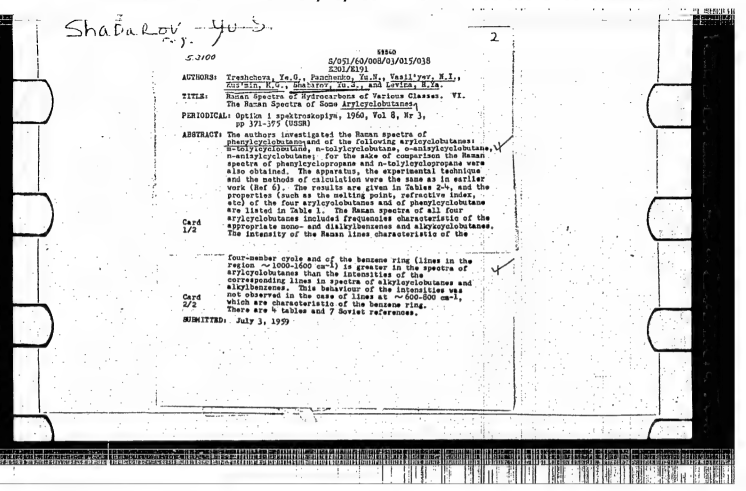
Card 3/4

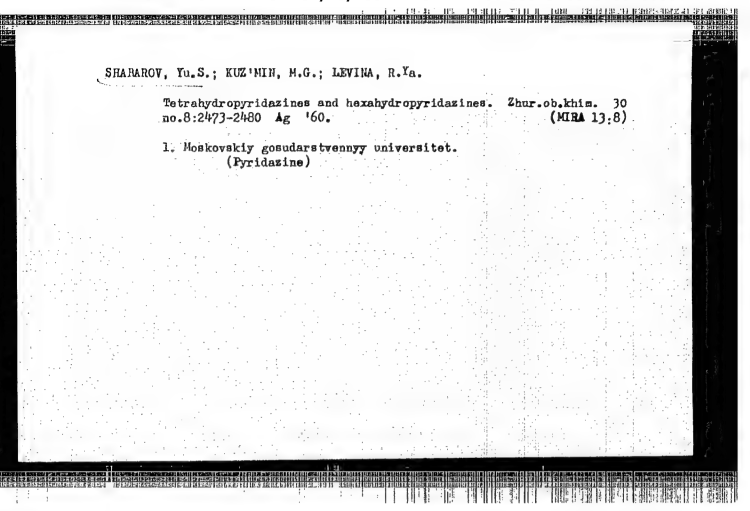
Cyclopropanes and Cyclobutanes. On the Decomposition SOV/20-127-1-29/65 of Alkyl-tetrahydropyridazines

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

PRESENTED: February 18, 1959, by A. N. Nesmeyanov, Academician

SUBMITTED: February 14, 1959





84871

s/079/60/030/010/007/030 B001/B075

11.1210

AUTHORS:

Shabarov, Yu. S., Levina, R. Ya., Kuz'min, M. G.

Vasil'yev, N. I., and Damir, N. A.

TITLE:

Cyclopropenes and Cyclobutanes XI. Methylphenyl

Cyclobutanes

PERIODICAL:

Zhurnal obshchey khimii, 1960, Vol. 30, No. 10,

pp. 3210-3214

TEXT: In their investigation (Refs. 1-4) on the catalytic splitting of alkyl- and aryl tetrahydropyridazines, the authors could obtain only aryl cyclobutanes, but no alkyl cyclobutanes (Ref. 4). In the present work, the catalytic splitting of alkyl aryl tetrahydropyridazines has been attempted for the purpose of synthesizing alkyl aryl cyclobutanes. The authors proceeded from 5-methyl- and 6-methyl-3-phenyl tetrahydropyridazines which were obtained by hydrolyzing the adducts of the corresponding dienes with azodicarboxylic acid ester. These initial products proved to be unstable compounds, and were identified from their addition products to phenyl isothiocyanate. The presence of an NH group in the initial products was also proved (Ref. 3). The catalytic splitting of the Card 1/2

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Cyclopropanes and Cyclobutanes. XI. Methylphenyl 5/079/60/030/010/007/030 Cyclobutanes 7 B001/B075

two initial pyridazines was carried out in the same way as that of aryltetrahydropyridazines described in Refs. 1-3. Thus, a gas composed of nitrogen and propylene evolved under the formation of a mixture of the corresponding methyl-phenyl cyclobutane and styrene. Consequently, the decomposition of alkyl aryl tetrahydropyridazines is similar to that of aryl- Δ^2 -tetrahydropyridazines: While nitrogen is produced, the biradical is formed as an intermediate which yields cyclobutane and two ethylene hydrocarbons. The molar ratio between the yield of methyl-phenyl cyclobutane and styrene proved to depend on the position of the methyl group in the tetrahydropyridazine ring. Vacuum distillation (cf. experimental part) of the resulting cyclobutane showed that 1-methyl-2-phenyl cyclobutane consisted of a mixture of cis- and trans-isomers (1:1), whereas 1-methyl-3-phenyl cyclobutane contained 80% of the trans-form. Raman spectra showed the absence of olefins in the separated hydrocarbons (Ref. 5). The characteristic frequencies 912-950 cm⁻¹ confirmed the presence of a four-membered ring. There are 1 table and 12 references: 9 Soviet, 2 German, and 1 French.

ASSOCIATION:

Moskovskiy gosudarstvennyy universitet

(Moscow State University)

SUBMITTED:

December 7, 1959

Card 2/2

"APPROVED FOR RELEASE: 07/20/2001

87525

s/079/60/030/012/005/027 B001/3064

5 5500

AUTHORS:

Shabarov, Yu. S., Levina, R. Ya., Potapov, V. K.,

Osipov, A. M., and Treshchova, Ye. G.

TTTLE:

Cyclopropanes and Cyclobutanes. XIV. Phenyl Cyclopropanes With Substituents in the Para Position of the Benzene Cycle

PERIODICAL:

Zhurnal obshchey khimii, 1960, Vol. 30, No. 12,

pp. 3874-3876

TEXT: In previous papers (Refs. 1-4) the authors reported on the effect of the nature of aryl radicals upon the reactivity of the three-membered cycle linked with it. Thus, it was found that polymerizability strongly increases under the action of AlCl, in the following order: phenyl cyclopropane < p-tolyl cyclopropane < p-anisyl cyclopropane (Ref. 2). The three-membered cycle which opens readily in phenyl cyclopropane (Ref. 3), p-tolyl cyclopropane, p-anisyl cyclopropane (Ref. 4) under the action of mercury salts, is stable in p-nitrophenyl cyclopropane (Ref. 1). To render these observations more complete, it is necessary to develop a method of synthesizing phenyl cyclopropane with various substituents in the benzene cycle. p-aminophenyl cyclopropane which could be easily obtained by Card 1/3

87525

Cyclopropanes and Cyclobutanes. XIV. Phenyl S/079/60/030/012/005/027 Cyclopropanes With Substituents in the Para B001/B064 Position of the Benzene Cycle

nitrating phenyl cyclopropane, with subsequent reduction of the nitro group to the amino group (Ref. 1), served as the initial product. The replacement of the latter in p-amino phenyl cyclopropane by other substituents was carried out by diazotization. Thus, p-hydroxy-p-chloro- and p-bromophenyl cyclopropane results: CH2

A study of the Raman spectra of the phenyl cyclopropanes obtained showed that no unsaturated compounds had been added; intensive frequencies appeared at 1600 cm⁻¹, which are characteristic of the aromatic cycle, as well as bands (1200-1260 cm⁻¹) indicating the presence of the phenyl cyclopropane molecule (Refs. 5, 6). The ultraviolet absorption curves of aryl cyclopropanes (Diagrams 1 and 2) showed the same character as those of p-tolyl Card 2/3

"APPROVED FOR RELEASE: 07/20/2001

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Cyclopropanes and Cyclobutanes. XIV. Phenyl Cyclopropanes With Substituents in the Para Position of the Benzene Cycle

S/079/60/030/012/005/027 B001/B064

cyclopropane (Ref. 5) and p-aminophenyl cyclopropane (Ref. 1). There are 2 figures and 6 references: 5 Soviet and 1 French.

ASSOCIATION:

Moskovskiy gosudarstvennyy universitet

(Moscow State University)

SUBMITTED:

January 14, 1960

Card 3/3

CIA-RDP86-00513R001548510006-1" APPROVED FOR RELEASE: 07/20/2001

"APPROVED FOR RELEASE: 07/20/2001

CIA-RDP86-00513R001548510006-1

5.3610

69997

AUTHORS:

Levina, L. Ya., Shabarov, Yu. S., Kuz'min, M. G.

S/020/60/131/05/027/069

B011/B117

TITLE:

On the Interaction Between Azodicarboxylic Esters and hem-Dialkyl

Butadienes

PERIODICAL: Doklady Akademii nauk SSSR, 1960, Vol 131, Nr 5, pp 1080-1083 (USSR)

TEXT: The authors continued their investigations of the decomposition of alkyl tetrahydropyridazine (Ref 1), and made an attempt to synthetize hem-dialkyl tetrahydropyridazines from the adducts of the azodicarboxylic ester with hem-dialkyl butadienes. They could establish for the first time that 1,1-dialkyl butadienes react with the azodicarboxylic ester through mobile hydrogen in the a-position (by substitution addition and not by diene synthesis), and not through the double bonds. It could be further proved by the authors that the adducts of the substances mentioned in the title have no cyclic structure (being no tetrahydropyridazine derivatives), but they are, in reality, monosubstituted hydrazo dicarboxylic esters (see schemes). The structure of the adduct (I) of 2,4-dimethyl pentadiene with an azodicarboxylic ester was established by means of its conversions: when subjected to cold hydrogenation, (I) adds two moles of hydrogen per one mole of (I), and, thus, contains two double bonds in the molecule. Hydrolysis of (I) yields a monosubstituted hydrazine (III). The

Card 1/3

69998

On the Interaction Between Azodicarboxylic Esters and hem-Dialkyl Butadienes

S/020/60/131/05/027/069 B011/B117

latter contains one NH2 group which was detected by obtaining the benzylidene derivative (IV) from (III). If the azodicarboxylic ester is reacted with 2,4-dimethyl pentadiene-1,3, two adducts (Va) and (Vb) (in reality their mixture) can be formed, since the mentioned pentadiene contains two types of methyl groups. After hydrogenation, however, both (Va) and (Vb) yield the same substituted hydrodicarboxylic ester (VI). By hydrolysis of (VI), the monosubstituted hydrazine (VII) is formed from which the benzylidene derivative (VIII) can be obtained. In addition, nitrogen is evolved under the action of mercuric oxide, with (VII) being converted to 2,4-dimethyl pentane (which is a reaction characteristic of monoalkyl hydrazines, reference 4). All these reactions are clearly indicative of the acyclic structure of the corresponding adducts. Obviously, the dienes used in this case react with the azodicarboxylic ester through a single methyl group only. An analogous reaction between maleic anhydride and olefines (Ref 8) takes place only at 200-250°, while the much more active azodicarboxylic ester reacts already at 20-50°. There are 9 references, 3 of which are Soviet.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

Card 2/3

On the Interaction Between Azodicarboxylic Esters
and hem-Dialkyl Butadienes

PRESENTED: December 14, 1959, by A. N. Nesmeyanov, Academician

SUBMITTED: December 12, 1959

Card 3/3

SHABAHOV, Yu.S.; VASIL'YEV, N.I.; MAMAYEVA, N.K.; LEVINA, R.Ya.

Hew method of synthesizing 3-aryl-1,4,5,6-tetrahydropyridazines.

\$\int_{-}^{0}\$-Diphenylylcyclobutane. Dokl. AN SSSR 135 no.4:879-882 '60.

(MIRA 13:11)

1. Hoskovskiy gosudarstvennyy universitet im. M.V.Louonosova.

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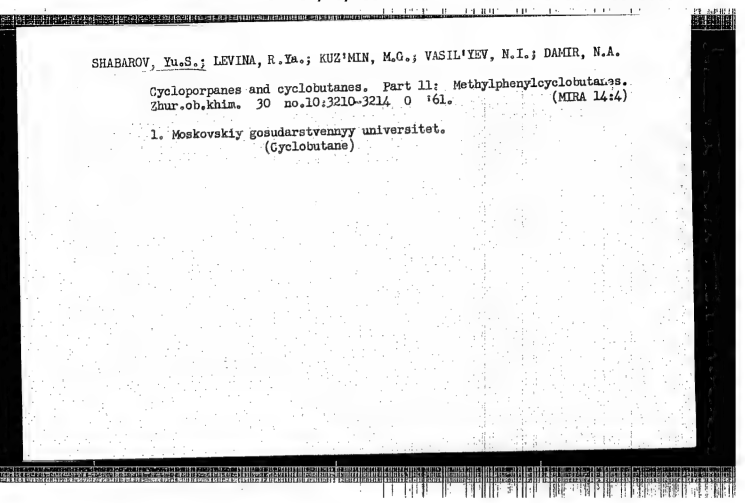
(Cyclobutane)

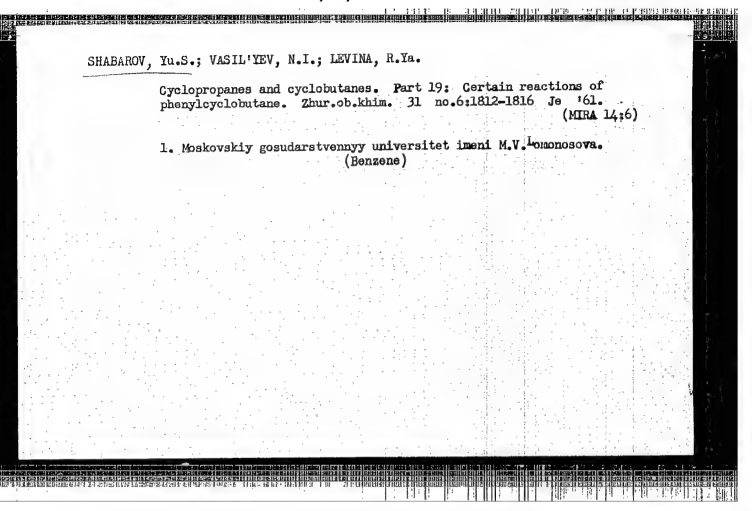
SHABAROV, Yu.S.; LEVINA, R.Ya.; VISIL'YEV, N.I.; VASILENKO, N.A.

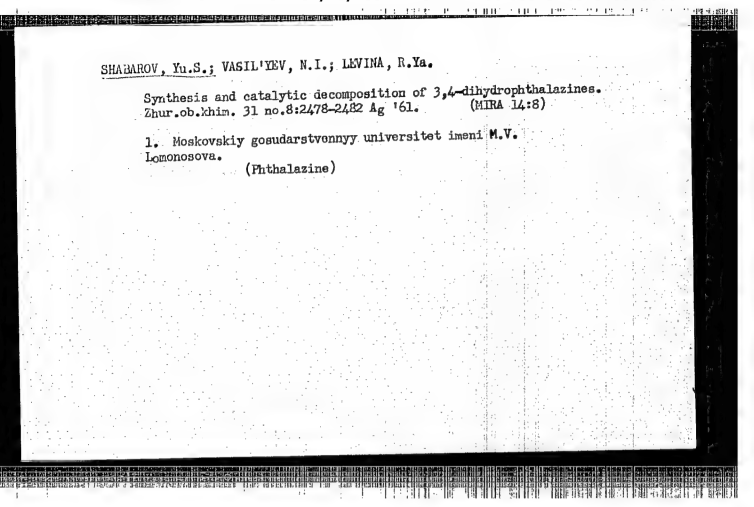
Cyclopropunes and cyclobutanes. Part 16: \(\rho \) -ohloro- and \(\rho \) -bromophenylcyclobutanes. Zhur. ob. khim. 31 no. 2:378-382

F '161. (KIKA 14:2)

1. Moskovskiy gosudarstvennyy universitet. (Gyclopropane) (Cyclobutane)







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KUTHORS:

Shchetinin, A. A., Topchiyeva, I. N., Shabarov, Yu. S.,

Levina, R. Ya.

TITLE:

Cyclopropanes and cyclobutanes. XXIII. Polymerization of

aryl cyclopropanes at high pressures and temperatures

PERIODICAL: Vysokomolekulyarnyye soyedineniya, v. 4, no. 4, 1962, 499-502

TEXT: The effect of high pressures (500-7500 atm) on phenyl-, p-aminophenyl-, p-dimethyl aminophenyl-, p-chlorophenyl, p-hydroxyphenyl-, p-methoxyphenyl cyclopropane, and p-methoxyphenyl cycloputane between

100 and 200°C was investigated. Of these only p-hydroxy- and p-methoxyphenyl cyclopropane undergo polymerization. The latter gives a

viscous oil at 500 atm and 200°C, a solid polymer at 7500 atm and 200°C. With hydroquinone as inhibitor only a viscous liquid is formed at 7500 atm. Therefore, polymerization takes place by the free-radical mechanism. Only cyclopropane rings participate in the polymerization. The end product of the thermal decomposition and the subsequent oxidation of the Card 1/2

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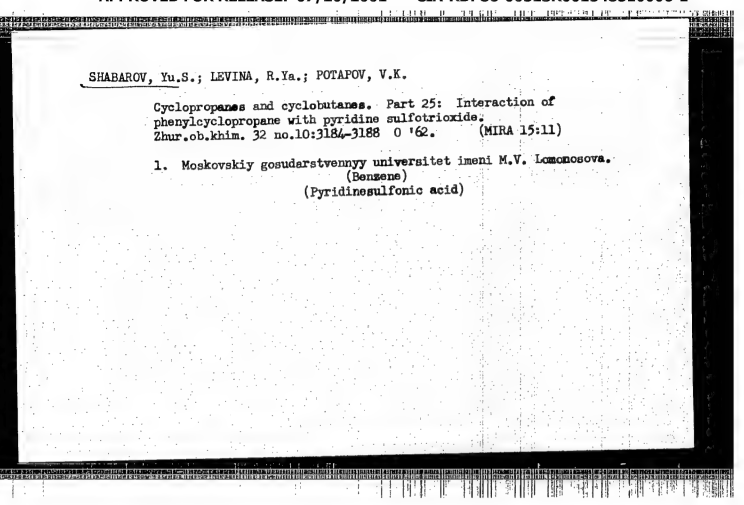
S/190/62/004/004/004/019
Cyclopropand and cyclobutanes. ... B119/B138

polymer is anisic acid. There is 1 table.

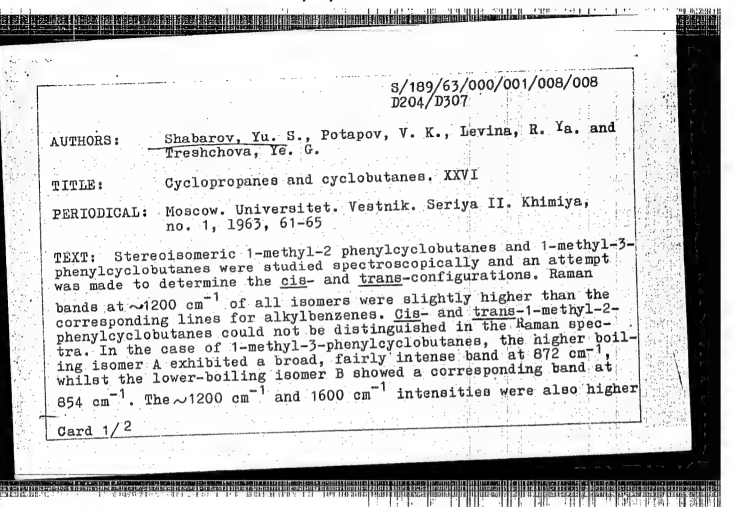
ASSOCIATION: Moskovskiy gosudarstvennyy universitet im. M. V. Lomonosova (Moscow State University imeni M. V. Lomonosov)

SUBMITTED: March 6, 1961

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S/189/63/000/001/008/008 Cyclopropanes and cyclobutanes ... D204/D307

in B, a property characteristic of trans-forms. Uv spectra of 1-methyl-2-phenylcyclobutanes in iso-octane showed that isomer A, distinguished by higher physical constants, absorbed more strongly in the 225 - 250 mu region than the other isomer, B. The same was true of the A-form of 1-methyl-3-phenylcyclobutane, though to a lesser extent. Control tests with 1,2-diphenylcyclopropanes, whose configurations were established chemically, showed that transforms absorbed more strongly in the uv. Configurations of 1-methyl-2-phenylcyclobutanes may thus be assigned only from uv absorption spectra, whilst the configurations of 1-methyl-3-phenyl-cyclobutanes remain unresolved, since Raman and uv spectra gave contradictory indications. L. A. Kazitsyna measured the uv absorption. There are 3 figures and 1 table.

ASSOCIATION: Kafedra organicheskoy khimii (Department of Organic

Chemistry)

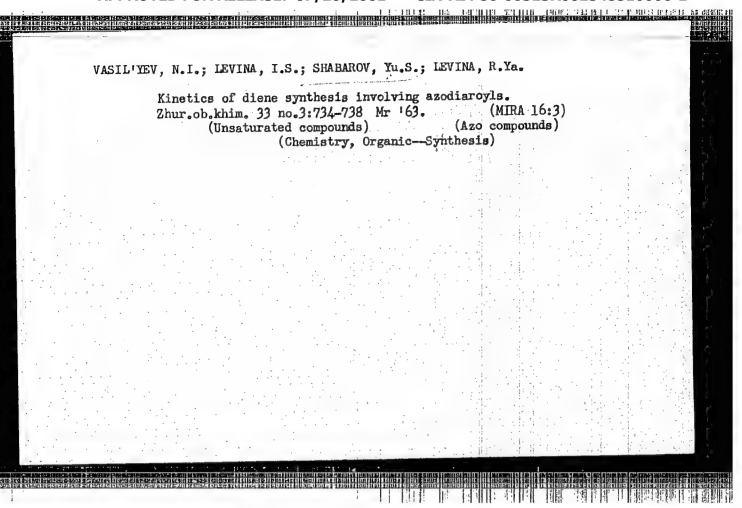
SUBMITTED: June 28, 1961

Card 2/2

SHABAROV, Yu.S.; POTAPOV, V.K.; LEVINA, R.Ya.; TRESHCHOVA, Ye.G.

Cyclopropanes and cyclobutanes. Part 26: Stereoisomeric
1,2- and 1,3-methylphenylcyclobutanes. Vest: Mosk.un. Ser.2: Khim.
18 no.1:61-65 Ja-F '63.

1. Kafedra organicheskoy khimii Moskovskogo universiteta.
(Cyclobutane) (Isomers)

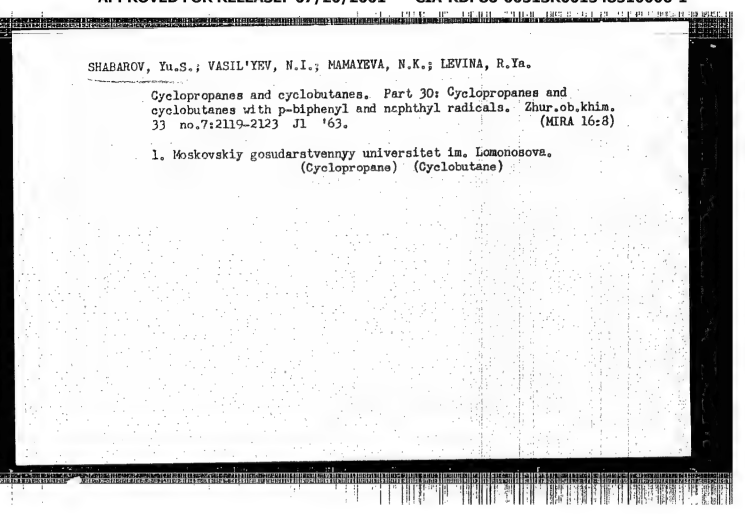


SHABAROV, Yu.S.; V.SIL'YEV, N.I.; MAMAYEVA, N.K.; LEVINA, R.Ya.

Reduction of pyridazinones and phthalazones by lithium aluminum hydride.

Zhur.ob.khim. 33 no.4:1206-1210 Ap '63. (MTRA 16:5)

1. Moskovskiy gosudarstvennyy universitet imeni M.V.Lomonosova. (Pyridazinone) (Phthalazinone) (Lithium aluminum hydride)



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